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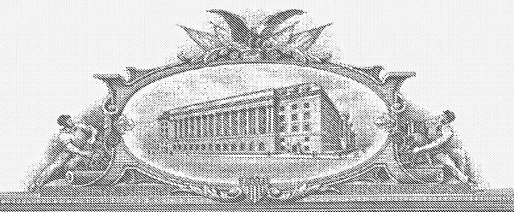
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APPLICATION NUMBER: 60/525,430
FILING DATE: November 26, 2003
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PROVISIONAL APPLICATION FOR PATENT COVER SHEET

This is a request for filing a PROVISIONAL APPLICATION FOR PATENT under 27 OPP (1997).

Express Mail Label No.

INVENTOR(S)									
Given Name (first and mid	ddle [if any])	Family Name or Surname		(City a	Residence (City and either State or Foreign Country)				
Congxin		Liang		Sunnyvale	Sunnyvale, California				
Additional inventors are being named on the0separately numbered sheets at						ereto			
TITLE OF THE INVENTION (500 characters max)									
Hydroxy Carboxy Pyrrolyl-indolinone Derivatives as Protein Kinase Inhibitors									
Direct all correspondence to: CORRESPONDENCE ADDRESS  Customer Number:									
OR									
Firm or Individual Name									
Address	Address 729 West Remington Drive								
Address						*			
City	Sunnyvale		State	CA	Zip	94087			
Country	USA		Telephone	408-746-0486	Fax	408-746-0486			
ENCLOSED APPLICATION PARTS (check all that apply)									
✓ Specification Number of Pages     10     CD(s), Number       ☐ Drawing(s) Number of Sheets     ✓ Other (specify)     Cover letter       ☐ Application Date Sheet. See 37 CFR 1.76						•			
		OR THIS PROVISIONAL AP	PLICATION FOR	PATENT					
Applicant claims small entity status. See 37 CFR 1.27.  A check or money order is enclosed to cover the filing fees.  The Director is herby authorized to charge filing fees or credit any overpayment to Deposit Account Number:  Payment by credit card. Form PTO-2038 is attached.					FILING FEE Amount (\$) \$80.00				
The invention was made by an agency of the United States Government or under a contract with an agency of the United States Government.  No.  Yes, the name of the U.S. Government agency and the Government contract number are:									
Respectfully submitted, SIGNATURE Congxin Liang TYPED or PRINTED NAME Congxin Liang TELEPHONE 408-746-0486				Date NUV, 25, 2003  REGISTRATION NO					

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FEE TO A NOBSITTAL	Complete if Known			
FEE TRANSMITTAL	Application Number			
for EV 2004	Filing Date	NOV. 25, 2003		
for FY 2004	First Named Inventor	CONGXIN LIANG		
ffective 10/01/2003. Patent fees are subject to annual revision.	Examiner Name			
plicant claims small entity status. See 37 CFR 1.27	Art Unit			

(\$) 80 00 TOTAL AMOUNT OF PAYMENT Attorney Docket No FEE CALCULATION (continued) METHOD OF PAYMENT (check all that apply) 3. ADDITIONAL FEES Money Other None Credit card Order Large Entity , Small Entity Deposit Account: 60 Fee Description Fee Paid Code (\$) Code (\$) Deposit Account 65 Surcharge - late filing fee or oath 2051 1051 130 Number Surcharge - late provisional filing fee or Deposit 1052 50 2052 Account cover sheet Name Non-English specification 1053 130 1053 130 e Director is authorized to: (check all that apply) 1812 2,520 For filing a request for ex parte reexamination 1812 2,520 Credit any overpayments Charge fee(s) indicated below Requesting publication of SIR prior to 920 1804 9201 1804 Charge any additional fee(s) or any underpayment of fee(s) Examiner action Requesting publication of SIR after Charge fee(s) indicated below, except for the filing fee 1805 1.840 1805 1.840\* Examiner action to the above-identified deposit account. 55 Extension for reply within first month 1251 110 2251 **FEE CALCULATION** Extension for reply within second month 1252 420 2252 210 1. BASIC FILING FEE 1253 950 2253 Extension for reply within third month arge Entity Small Entity Fee Paid Fee Description Extension for reply within fourth month Fee Fee Code (\$) 1254 1.480 2254 740 1.005 Extension for reply within fifth month 2255 1255 2.010 1001 770 2001 385 Utility filing fee 1401 330 2401 165 Notice of Appeal 1002 340 2002 170 Design filing fee 165 Filing a brief in support of an appeal 330 2402 1402 2003 265 Plant filing fee 1003 530 145 Request for oral hearing 1403 290 2403 1004 770 2004 385 Reissue filing fee 80.00 1451 1,510 1451 1,510 Petition to institute a public use proceeding 2005 (80) Provisional filing fee 1005 160 55 Petition to revive - unavoidable 1452 110 2452 SUBTOTAL (1) (\$) \$12 1453 1,330 2453 665 Petition to revive - unintentional 2. EXTRA CLAIM FEES FOR UTILITY AND REISSUE 2501 665 Utility issue fee (or reissue) 1501 1.330 Fee from Fee Paid Extra Claims below 1502 480 2502 240 Design issue fee Total Claims -20 1503 640 2503 320 Plant issue fee Independent 130 Petitions to the Commissioner 1460 130 1460 Multiple Dependent 50 Processing fee under 37 CFR 1.17(q) 1207 50 1807 Large Entity 180 Submission of Information Disclosure Stmt Small Entity 180 1806 1806 Fee Fee Code (\$) Fee Description 40 Recording each patent assignment per Code (\$) 8021 40 8021 property (times number of properties) Claims in excess of 20 1202 2202 9 18 385 Filing a submission after final rejection (37 CFR 1.129(a)) 1809 770 2809 Independent claims in excess of 3 1201 86 2201 43 385 For each additional invention to be 1203 290 2203 145 Multiple dependent claim, if not paid 1810 770 2810 examined (37 CFR 1.129(b)) Reissue independent claims 1204 86 2204 43 385 Request for Continued Examination (RCE) 2801 over original patent 1801 770 900 Request for expedited examination 1802 900 1802 \*\* Reissue claims in excess of 20 1205 18 2205 of a design application and over original patent Other fee (specify) (\$) SUBTOTAL (2) \*Reduced by Basic Filing Fee Paid (\$) SUBTOTAL (3) or number previously paid, if greater; For Reissues, see above

SUBMITTED BY				(Complete (if applicable))		
Name (Print/Type)	CONGXIN LIANG	Registration No. (Attorney/Agent)	Telephone	408-746-0486		
Signature	Cinji 2 mus		Date	Nov. 25, 2003		

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Nov. 25, 2003

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Mail Stop Provisional Patent Application Commissioner for Patents Box 1450 Alexandria, VA 22313-1450

Dear Sir or Madam:

Enclosed please find the following documents for a provisional patent application:

- Provisional Application for Patent Cover Sheet
- Fee transmittal for FY 2004

Coryi Dinny

- Credit card payment form (for \$80.00)
- Description of the invention: Hydroxy Carboxy Pyrrolyl-indolinone Derivatives as Protein Kinase Inhibitors (10 pages)

Please check the list and call me at (408)-718-9689 (mobile) if the application is incomplete.

Best regards,

Congxin Liang

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## HYDROXY CARBOXY PYRROLYL-INDOLINONE DERIVATIVES AS PROTEIN KINASE INHIBITORS

#### **BACKGROUND OF THE INVENTION**

#### Field of Invention

This invention relates to certain hydroxy carboxy pyrrolyl-indolinone derivatives and their pharmaceutically acceptable salts as protein kinase inhibitors. The compounds of this invention are therefore useful in treating disorders related to abnormal protein kinase activities such as cancer.

#### State of the Art

Protein kinases are enzymes that catalyze the phosphorylation of hydroxyl groups of tyrosine, serine, and threonine residues of proteins. Many aspects of cell life (for example, cell growth, differentiation, proliferation, cell cycle and survival) depend on protein kinase activities. Furthermore, abnormal protein kinase activity has been related to a host of disorders such as cancer and inflammation. Therefore, there is a great deal of effort directed to identifying ways to modulate protein kinase activities. In particular, many attempts have been made to identify small molecules which act as protein kinase inhibitors.

#### **DESCRIPTION OF THE INVENTION**

This invention discloses that certain hydroxy carboxy pyrrolyl-indolinone derivatives may have interesting and unexpected properties that advantageously distinguish them from known compounds. They are therefore useful in treating disorders related to abnormal protein kinase activities such as cancer.

One embodiment of this invention is a compound of Formula (I) or (II):

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(I) 
$$R^{3} \longrightarrow NR^{5}-(CHR^{6})_{n}-CH(OH)-(CHR^{7})_{m}-COOR^{8}$$

$$R^{1} \longrightarrow NH$$

(II)

$$R^3$$
 $NR^5$ -(CHR6)<sub>n</sub>-(CH(OH)-CH<sub>2</sub>)<sub>m</sub>-COOR8

 $R^4$ 

wherein:

R<sup>1</sup> is selected from the group consisting of hydrogen, halo, alkyl, cycloalkyl, haloalkyl, hydroxy, alkoxy, amino, alkylamino, amide;

R<sup>2</sup> is selected from the group consisting of hydrogen, halo, alkyl, cycloalkyl, haloalkyl, hydroxy, alkoxy, amino, alkylamino, amide, aryl, heteroaryl, sulfonyl, sulfonamide:

R<sup>3</sup> is selected from the group consisting of hydrogen, alkyl, aryl, heteroaryl, and amide;

R<sup>4</sup> is selected from the group consisting of hydrogen, alkyl;

R<sup>5</sup> and R<sup>6</sup> are independently hydrogen or alkyl;

R<sup>7</sup> is hydrogen, alkyl or hydroxyl;

R<sup>8</sup> is selected from the group consisting of hydrogen, alkyl, cycloalkyl, substituted alkyl;

n and m are independently 0, 1, 2, or 3; or a pharmaceutically acceptable salt, prodrug thereof. It may also act as a prodrug. The compound of Formula (I) or (II) may exist in or co-exist with its cyclic lactone form in

solution or in vivo.

Another embodiment of this invention is a compound of Formula (Ia) or (IIa):

(Ia) 
$$R^3$$
 $NR^5$ -(CHR $^6$ )<sub>n</sub>-CH(OH)-(CHR $^7$ )<sub>m</sub>-COOR $^8$ 
 $R^4$ 

(IIa) 
$$R^3$$
  $NR^5$ -(CHR6)<sub>n</sub>-(CH(OH)-CH<sub>2</sub>)<sub>m</sub>-COOR8  $R^4$ 

wherein:

R<sup>1</sup> and R<sup>2</sup> are independently hydrogen or fluoro and are located, respectively, at the 4- or 5-position of the indolinone ring;

R<sup>3</sup> and R<sup>4</sup> are methyl;

R<sup>5</sup>, R<sup>6</sup>, and R<sup>8</sup> are hydrogen;

R<sup>7</sup> is hydrogen or hydroxyl;

n and m are independently 1, or 2;

or a pharmaceutically acceptable salt thereof. It may also act as a prodrug. The compound of Formula (Ia) or (IIa) may exist in or co-exist with its cyclic lactone form in solution or in vivo.

Another embodiment of this invention is a compound of Formula (IIb):

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(IIb) 
$$R^3$$
  $NR^5$ -(CHR<sup>6</sup>)<sub>n</sub>-(\*CH(OH)-CH<sub>2</sub>)<sub>m</sub>-COOR<sup>8</sup>

wherein:

 $R^1$  and  $R^2$  are independently hydrogen or fluoro and are located, respectively, at the 4- or 5-position of the indolinone ring;

R<sup>3</sup> and R<sup>4</sup> are methyl;

R<sup>5</sup>, R<sup>6</sup>, and R<sup>8</sup> are hydrogen;

n and m are 2;

or a pharmaceutically acceptable salt thereof. It may also act as a prodrug. Preferably, the stereochemistry at the  ${}^*C$  is (R).

The compound of Formula (IIb) may exist in or co-exist with its cyclic lactone form with Formula (IIc) in solution or *in vivo*:

(IIc) 
$$\mathbb{R}^2$$
  $\mathbb{R}^4$   $\mathbb{R}^4$ 

wherein:

R<sup>1</sup> and R<sup>2</sup> are independently hydrogen or fluoro and are located, respectively, at the 4- or 5-position of the indolinone ring;

 $R^3$  and  $R^4$  are methyl.

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Representative compounds of the present invention are shown below.

In the above examples, 1a is the cyclic lactone of 1 and they may co-exist in solution or *in vivo*. Similarly, 2a is the cyclic lactone of 2 and they may co-exist in solution or *in vivo*. Furthermore, in the above examples the stereochemistry at the carbon atom carrying a hydroxyl group is either RS, R, or S. In 1, 1a, 2, and 2a, such stereochemistry is preferably R.

#### Utility

The present invention provides compounds capable of regulating and/or modulating protein kinase activities of, but not limited to, VEGFR and/or PDGFR. Thus, the present invention provides a therapeutic approach to the treatment of disorders related to the abnormal functioning of these kinases. Such disorders include, but not limited to,

#### Provisional Patent Application, Congxin Liang, 11/26/2003, Page 6 of 10

solid tumors such as glioblastoma, melanoma, and Kaposi's sarcoma, and ovarian, lung, prostate, pancreatic, colon and epidermoid carcinoma. In addition, VEGFR/PDGFR inhibitors may also be used in the treatment of restenosis and diabetic retinopathy.

Furthermore, this invention relates to the inhibition of vasculogenesis and angiogenesis by receptor-mediated pathways, including the pathways comprising VEGF receptors, and/or PDGF receptors. Thus the present invention provides therapeutic approaches to the treatment of cancer and other diseases which involve the uncontrolled formation of blood vessels.

#### **Synthesis of Compounds**

The compounds of this invention can be synthesized by following the published general procedures (e.g. Sun et al., 2003, J. Med. Chem., 46:1116-119). But the following intermediates are specific to compounds of this invention and may be used in place of their respective counterparts in the above-mentioned general procedure: 4,5-difluoro-oxindole; (4R,6R)-t-butyl-6-(2-aminoethyl)-2,2-dimethyl-1,3-dioxane-4-acetate; and 4-amino-3-hydroxy-butanic acid. These intermediates may be purchased from commercial sources (e.g. Fisher Scientific, Fairlawn, New Jersey). Another variation from the above-mentioned general procedure is that in the synthesis of 1/1a and 2/2a using (4R,6R)-t-butyl-6-(2-aminoethyl)-2,2-dimethyl-1,3-dioxane-4-acetate, the protecting groups need to be removed from the final product. Yet another variation from the above-mentioned general procedure is that in the synthesis of 3 and 4 using 4-amino-3-hydroxy-butanic acid, the acid needs to be protected before amidation and the protection group needs to be removed from the final product. These variations from the above-mentioned general procedure can be understood and carried out by those skilled in the art. Thus, the compounds of the present invention can be synthesized by those skilled in the art.

The compounds described herein are presently representative of preferred embodiments, are exemplary, and are not intended as limitations on the scope of the invention. It will be readily apparent to one skilled in the art that varying substitutions and modifications may be made to the invention disclosed herein without departing from the scope and spirit of the invention.

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#### The Claims

What is claimed is:

1. A compound of Formula (I):

(I) 
$$R^3$$
 $NR^5$ -(CHR $^6$ )<sub>n</sub>-CH(OH)-(CHR $^7$ )<sub>m</sub>-COOR $^8$ 
 $R^1$ 
 $R^4$ 

wherein:

R<sup>1</sup> is selected from the group consisting of hydrogen, halo, alkyl, cycloalkyl, haloalkyl, hydroxy, alkoxy, amino, alkylamino, amide;

R<sup>2</sup> is selected from the group consisting of hydrogen, halo, alkyl, cycloalkyl, haloalkyl, hydroxy, alkoxy, amino, alkylamino, amide, aryl, heteroaryl, sulfonyl, sulfonamide;

R<sup>3</sup> is selected from the group consisting of hydrogen, alkyl, aryl, heteroaryl, and amide;

R<sup>4</sup> is selected from the group consisting of hydrogen, alkyl;

 ${\bf R}^5$  and  ${\bf R}^6$  are independently hydrogen or alkyl;

R<sup>7</sup> is hydrogen, alkyl, or hydroxyl;

R<sup>8</sup> is selected from the group consisting of hydrogen, alkyl, cycloalkyl, substituted alkyl;

n and m are independently 0, 1, 2, or 3; or a pharmaceutically acceptable salt, prodrug thereof.

2. A compound of Formula (II):

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wherein:

R<sup>1</sup> is selected from the group consisting of hydrogen, halo, alkyl, cycloalkyl, haloalkyl, hydroxy, alkoxy, amino, alkylamino, amide;

R<sup>2</sup> is selected from the group consisting of hydrogen, halo, alkyl, cycloalkyl, haloalkyl, hydroxy, alkoxy, amino, alkylamino, amide, aryl, heteroaryl, sulfonyl, sulfonamide;

R<sup>3</sup> is selected from the group consisting of hydrogen, alkyl, aryl, heteroaryl, and amide;

R<sup>4</sup> is selected from the group consisting of hydrogen, alkyl;

R<sup>5</sup> and R<sup>6</sup> are independently hydrogen or alkyl;

R<sup>8</sup> is selected from the group consisting of hydrogen, alkyl, cycloalkyl, substituted alkyl;

n and m are independently 0, 1, 2, or 3; or a pharmaceutically acceptable salt, prodrug thereof.

#### 3. The compound of claim 1, wherein:

R<sup>1</sup> and R<sup>2</sup> are independently hydrogen or fluoro and are located, respectively, at the 4- or 5-position of the indolinone ring;

R³ and R⁴ are methyl;

R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> are hydrogen;

n and m are independently 0, 1, or 2;

or a pharmaceutically acceptable salt thereof.

#### 4. The compound of claim 2, wherein:

 $R^1$  and  $R^2$  are independently hydrogen or fluoro and are located, respectively, at the 4- or 5-position of the indolinone ring;

R<sup>3</sup> and R<sup>4</sup> are methyl;

### Provisional Patent Application, C ngxin Liang, 11/26/2003, Page 9 of 10

R<sup>5</sup>, R<sup>6</sup>, and R<sup>8</sup> are hydrogen;

n and m are independently 1, or 2;

or a pharmaceutically acceptable salt thereof.

#### 5. A compound of Formula (IIb):

(IIb) 
$$R^3$$
  $NR^5$ -(CHR<sup>6</sup>)<sub>n</sub>-(\*CH(OH)-CH<sub>2</sub>)<sub>m</sub>-COOR<sup>8</sup>

wherein:

 $R^1$  and  $R^2$  are independently hydrogen or fluoro and are located, respectively, at the 4- or 5-position of the indolinone ring;

R<sup>3</sup> and R<sup>4</sup> are methyl;

R<sup>5</sup>, R<sup>6</sup>, and R<sup>8</sup> are hydrogen;

n and m are 2;

or a pharmaceutically acceptable salt thereof.

#### 6. A compound of Formula (IIc):

(IIc) 
$$\mathbb{R}^2$$
  $\mathbb{N}$   $\mathbb{R}^4$ 

wherein:

R<sup>1</sup> and R<sup>2</sup> are independently hydrogen or fluoro and are located, respectively, at the 4- or 5-position of the indolinone ring;

R<sup>3</sup> and R<sup>4</sup> are methyl.

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7. The compound or salt of claim 2, wherein the compound is selected from the group consisting of:

8. The compound or salt of claim 1, wherein the compound is selected from the group consisting of:

- 9. A method for the modulation of the catalytic activity of a protein kinase with a compound or salt of any one of claims 1, 2, 3, 4, 5, 6, 7, or 8.
- 10. The method of claim 9, wherein said protein kinase is selected from the group consisting of VEGF receptors, PDGF receptors.